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The Dinegentropy of Diagnostic and Detection Tests

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1 Introduction.

Assessing the efficacy of diagnostic tests when their receiver operating characteristic curves cross has been an open problem for quite some time. In this note, we outline a proposed solution to this problem based on an information theoretic measure which I. J. Good has labeled *dinegentropy*. We illustrate the effectiveness of this measure, empirically, via an example. The detailed development of our arguments which lead to the proposed measure have been delegated to a more comprehensive document.

2 Preliminaries

Diagnostic testing plays a key role in medicine, threat detection, supervised classification, signal processing, the verification and validation of test ban treaties, alerts for tornadoes, tsunamis and earthquakes and the evaluation zero-net investment Strategies in finance. However, diagnostic tests are not perfect; they are prone to misdiagnosis and the generation of false alarms.

Assume a disease (or threat) we are interested in diagnosing (or detecting). The disease spawns a single measurement variable Z , that is compared to a threshold T . An individual is classified D - for diseased - if $Z > T$; otherwise the individual is classified N - for normal. The classification is done by a

diagnostician (who is a decision maker), say \mathcal{D} .

Were T be set to say t , then the *sensitivity* of the test, denoted $S_e(t)$ is the probability of correctly classifying a diseased individual. Similarly, the probability of correctly classifying a normal individual is denoted by $S_p(t)$, the *specificity* of the test. Ideally, one wants both $S_e(t)$ and $S_p(t)$ to be close to one, but for this to happen, the distribution of Z for D and N should have little or no overlap. Otherwise an increase in $S_e(t)$ will cause a decrease in $S_p(t)$ and vice-versa. The parameters $S_e(t)$ and $S_p(t)$ are therefore *adversarial*.

3 The Receiver Operating Characteristic Curve.

The adversarial character of sensitivity and specificity is encapsulated via the *receiver operating characteristic curve* (ROC) which is a plot of $(1 - S_p(t))$ versus $S_e(t)$. The ROC is monotonically increasing, continuous, and differentiable everywhere. It is concave in $(1 - S_p(t))$, if for all z , $F_0(z) \geq F_1(z)$; it will have a convex segment whenever $F_1(z) > F_0(z)$. Here, F_0 (F_1) are the distributions of Z under N (D). When F_0 and F_1 are specified, then two tests, each having their own classification thresholds can be compared to assess as to which of the two superior. An omnibus way to do so is via the area under the ROC, abbreviated AUC. The AUC is a satisfactory metric of comparison when the ROC's of the two tests do not cross. This will happen when F_0 and F_1 do not cross.

Note that the $AUC \in (0, 1)$, with 1 corresponding to a test providing a perfect diagnosis, and an $AUC = .5$ being given by a test whose diagnosis is random; i.e. diagnosis by the flip of a coin. All ROC's below the diagonal line correspond to tests that are worse than a random test.

The extent to which a diagnostic test is superior to a random diagnosis is

also reflected by what we introduce here as the *Gini Coefficient*, G_{F_0, F_1} , not to be confused with the Gini Index of economic inequality. Specifically,

$$G_{F_0, F_1} = 2(AUC) - 1. \quad (3.1)$$

$G_{F_0, F_1} = 1$, when the diagnostic is perfect; it is 0 when the test is random.

When the ROC's of two tests cross, alternatives to the AUC have been proposed. These are ad hoc, and their credibility remains to be assessed. In what follows we offer an approach to comparing two diagnostic tests over a range of applications, when their ROC's cross. Our approach is based on information theoretic considerations and is therefore grounded in this theory. The efficacy of our approach is illustrated by an example.

4 The Dinegentropy of an ROC Curve.

We propose the *Jeffreys-Good distance* or per Good (1987), the *dinegentropy* of an ROC as an omnibus measure for comparing two ROC curves, irrespective of their crossing properties. We motivate a use of this measure by considering the *Kullback-Leibler distance* between a reflected ROC curve, viewed as a distribution function, and the diagonal ROC curve as a reference distribution function, and then reversing the roles of these distributions to obtain a complementary Kullback-Leibler distance. The dinegentropy is the sum of the above two distance. This process leads to the following theorem; the detailed proof is in Singpurwalla (2018).

Theorem:

The dinegentropy of an ROC curve with $f_0(f_1)$ as the probability densities of $F_0(F_1)$ is:

$$\int_0^{\infty} [f_0(t) - f_1(t)] \log_2 \left[\frac{f_0(t)}{f_1(t)} \right] dt.$$

5 Demonstration of Efficacy.

The figure below demonstrates the efficacy of the dinegentropy measure for distinguishing between two diagnostic tests whose ROC's cross but which have the same value of the AUC. The $S_e(t)$'s of the two tests are based on the distribution functions of beta-distributed random variables with parameters 2 and 6, and 1 and 3, respectively. These distribution functions cross, but their resulting AUC's are the same, namely, .75. By contrast their dinegentropies differ, namely, 4.125 for the former, and 1.923 for the latter.

It has been argued in the reference mentioned above, that the diagnostic test with the larger dinegentropy is the better test.

6 References.

- [1] GOOD I. J. (1989). "The Contributions of Jeffreys to Bayesian Statistics". *Bayesian Analysis in Econometrics and Statistics: Essays in Honor of Harold Jeffreys*, ed. by A. Zellner. Kreiger Publishing. pp 21-34.
- [2] SINGPURWALLA, N. D. (2018). "The Stochastics of Diagnostic and Threat Detection Tests". Under Review.

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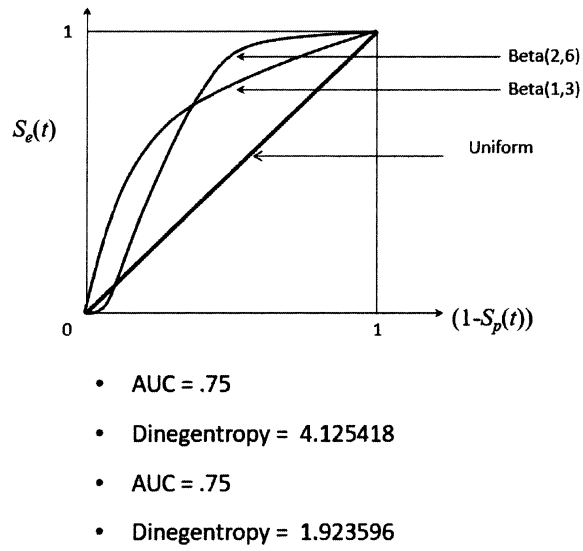


Figure 1: Demonstrating the Efficacy of Dinegentropy.